



Genomes & Developmental Control

The *cis*-regulatory system of the *tbrain* gene: Alternative use of multiple modules to promote skeletogenic expression in the sea urchin embryoMary E. Wahl^a, Julie Hahn^b, Kasia Gora^c, Eric H. Davidson^b, Paola Oliveri^{d,*}^a Department of Molecular and Cellular Biology, Harvard University, 16 Divinity Avenue, Cambridge, MA 02138, USA^b Division of Biology, California Institute of Technology, Pasadena, CA 91125, USA^c Department of Biology, Massachusetts Institute of Technology, 77 Massachusetts Avenue, Cambridge, MA 02139, USA^d Department of Genetics, Evolution and Environment, University College London, Gower Street, London WC1E 6BT, UK

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ABSTRACT

The genomic *cis*-regulatory systems controlling regulatory gene expression usually include multiple modules. The regulatory output of such systems at any given time depends on which module is directing the function of the basal transcription apparatus, and ultimately on the transcription factor inputs into that module. Here we examine regulation of the *Strongylocentrotus purpuratus tbrain* gene, a required activator of the skeletogenic specification state in the lineage descendant from the embryo micromeres. Alternate *cis*-regulatory modules were found to convey skeletogenic expression in reporter constructs. To determine their relative developmental functions in context, we made use of recombineered BAC constructs containing a GFP reporter and of derivatives from which specific modules had been deleted. The outputs of the various constructs were observed spatially by GFP fluorescence and quantitatively over time by QPCR. In the context of the complete genomic locus, early skeletogenic expression is controlled by an intron enhancer plus a proximal region containing a HesC site as predicted from network analysis. From ingresson onward, however, a dedicated distal module utilizing positive Ets1/2 inputs contributes to definitive expression in the skeletogenic mesenchyme. This module also mediates a newly discovered negative Erg input which excludes non-skeletogenic mesodermal expression.

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Introduction

The sea urchin regulatory gene *tbrain* (*tbr*) is zygotically expressed in the skeletogenic mesoderm (SM) of the cleavage and blastula stage embryo (Crocce et al., 2001; Oliveri et al., 2002), and its expression is required for the postgastrular formation of the larval spicules (Fuchikami et al., 2002). Through transcriptional activation of a target gene, *erg*, *tbr* establishes an *erg-hex-tgif-alx1* positive feedback circuit that maintains the regulatory state of the skeletogenic mesoderm domain from early in development and eventually, together with other regulators, serves as a transcriptional driver of an initial set of differentiation genes (Oliveri et al., 2008). The *tbr* gene thus has essential roles, first in specification of the SM and then in definitive larval skeletogenesis. Yet these roles, and the circuitry underlying them, are evolutionarily derived traits, since only modern sea urchins precociously segregate a SM lineage. In the sister group to the echinoids, the sea cucumbers, *tbr* is expressed in the developing endomesoderm (Maruyama, 2000). This is the pleisiomorphic function of the *tbr* gene in embryogenesis, since it is also expressed in endomesoderm in the more distant sea star outgroup (Hinman and

Davidson, 2007; Hinman et al., 2003; Shoguchi et al., 2000). Thus from an evolutionary standpoint the *tbr cis*-regulatory system is of particular interest since it must be at least partly “new”, and since it is a key mechanistic component of the skeletogenic micromere specification network: this, as a whole, is in itself a derived embryonic feature of the modern sea urchins (euechinoids).

Despite the simple pattern of *tbr* expression, which is confined entirely to the SM lineage throughout embryonic development, the *cis*-regulatory system of the *tbr* gene is anything but simple. Typically for regulatory genes (cf. Davidson, 2006), *tbr* is controlled by multiple *cis*-regulatory modules. Regulatory modules were identified in an intron as well as proximally in the closely related (actually congeneric) *strongylocentrotid* known as *Hemicentrotus pulcherrimus* (Ochiai et al., 2008). A different, also completely specific skeletogenic *cis*-regulatory module exists some distance upstream of the gene in *Strongylocentrotus purpuratus*, as we describe below. A major objective of this work was to resolve the various roles of these modules. Gene regulatory network analysis had shown that *tbr* lies under control of a double-negative gate (Oliveri et al., 2002, 2003, 2008; Revilla-i-Domingo et al., 2007). Thus the early zygotically expressed micromere repressor Pmar1 acts to prevent transcription in micromeres of the *hesC* gene, which encodes a dedicated repressor zygotically expressed everywhere in the embryo except in micromeres expressing the *pmar1* gene. Among the targets of HesC repres-

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sion is *tbr*, along with a small number of other initial founders of the SM regulatory state. The double-negative gate thus results in derepression of the *tbr* gene in the SM lineage. The putative site of HesC interaction in *tbr* cis-regulatory DNA had been identified (Ochiai et al., 2008), but there was little detailed information as to hesC effects on the *tbr* cis-regulatory system. In addition cis-regulatory mutations as well as other evidence indicated that some member(s) of the Ets family of transcription factors are required for *tbr* expression (Fuchikami et al., 2002; Ochiai et al., 2008). On the other hand, it had also been reported that morpholino-substituted antisense oligonucleotides (MASO) directed against the *S. purpuratus* Ets family members *SpErg*, *SpEts1/2*, and *SpTel* had no very significant effect (i.e., caused <3-fold change) on *tbr* expression up to 24 hpf (Oliveri et al., 2008). The role of Ets factors in *tbr* regulation altogether was clearly in need of further investigation. An additional mystery was that by late mesenchyme blastula stage *hesC* expression disappears from the non-skeletogenic mesenchyme (NSM; Smith and Davidson, 2008b), and *ets* expression spreads to include the NSM (Rizzo et al., 2006); yet *tbr* expression does not expand, remaining confined to the SM. Thus there appeared to be a need for either an additional yet unidentified NSM repressor of *tbr* expression, or a spatially-dedicated SM activator of *tbr* in later stages.

These issues are resolved in the cis-regulatory analyses described in this paper. The approach we have taken differs from the conventional in that we have attempted to examine cis-regulatory modular function in the context of the complete genomic *tbr* locus. To this end we utilized recombineered BAC reporters bearing module deletions or site mutations. Thus we have been able to establish the sequence of module deployment as well as determine the functionality of key transcription factor target sites. Perhaps not surprisingly, some of the insights we obtained as to module function in context proved invisible from the vantage point of the usual minimal expression constructs.

Materials and methods

BAC homologous recombination

Deletions of the $\gamma(2)$, B, and C modules from an *SpTbrain* GFP knock-in BAC (Damle et al., 2006) by homologous recombination were performed as described by Lee et al. (2007). The parental BAC is referred to as *tbr::GFP* BAC in the following. To produce a targeting cassette with homology to the regions bordering each module, a kanamycin resistance gene flanked by *frt* sites was amplified with the following primer pairs:

$\Delta\gamma(2)$ module-Forward: 5'-GACATAGGTATTTCTTATACATCGTCA-TGATTATGGTTACTCTCTAGATAACTGATCAGCTT-3'

$\Delta\gamma(2)$ module-Reverse: 5'-ATATATCTATAATTATATGGAATAA-TTCCATGAAATCTCATGTGGAGCTATTCCAGAAGTAGTA-3'

ΔB module-Forward: 5'-GGTAGTCACAAAGCCAAATACCTTACAA-GCTCTCTTTTATGTCGGAGTATCTAAGTACTCTTTGTAAAGCTGTC-TAATTTTCTGATTCTAGATAACTGATCAGCTT-3'

ΔB module-Reverse: 5'-AAATTCGTACGTTACTTTGAAATGAACCGAC-AATGCGGATTATAAGAGCTATTCCAGAAGTAGTA-3'

ΔC module-Forward: 5'-CAGCTTAGGCACTTTAACAAAAAAGAGTC-TTTAGAATCTTTGATCTAGATAACTGATCAGCTT-3'

ΔC module-Reverse: 5'-GAGCAAATCCTACATGATATCTACAGACA-TCATCAGATGCTTCAGGAGCTATTCCAGAAGTAGTA-3'.

Underlined sequences are homologous to the targeting cassette. Correct integration of the cassette into *tbr::GFP* BAC was confirmed by sequencing and diagnostic PCR. After removal of *kan^R* by induction of flippase, a 125-bp artifact of the cassette remained in the former location of each module.

To avoid undesired flippase recombination with a *frt* site at the GFP insertion site, mutation of the HesC binding site on *tbr::GFP* BAC was performed using a GalK positive/negative selection method (Warming et al., 2005). A targeting cassette containing *galK* was amplified with the following primers to introduce homology to the region flanking the HesC binding site:

HesCmut-cassette-Forward: 5'-CAGACTATTTTTCTTCTTCGTCGT-CGTCTAAATGTTATTTTCGAGTCGCCTGTTGACAATAATCATC-3'

HesCmut-cassette-Reverse: 5'-GGGCTACCAGACAATGGAGAGTCG-CGCGTTGATTGGCCGCCAGGGAGGTCAGCACTGTCTGCTCCTT-3'.

Underlined sequences are homologous to the targeting cassettes. After proper insertion was confirmed by sequencing and diagnostic PCR, the cassette was replaced with the mutated HesC site through homologous recombination with the following annealed oligonucleotides:

HesCmut-cassette-removal-Forward: 5'-CAGACTATTTTTCTTCT-TCTGTCGTCGTCTAAATGTTATTTTCGAGTCGGACTCTCCCTGGCGGCC-AATCAACGCGCGACTCTCGATTGTCTGGTAGCCC-3'

HesCmut-cassette-removal-Reverse: 5'-GGGCTACCAGACAATCGA-GAGTCGCGCGTTGATTGGCCGCCAGGGAGGAGTCCGACTCGAAA-TAACATTTAGACGACGACGAAGAAGAAAAAATAGTCTG-3'.

Generation of cis-regulatory reporter constructs

The $\gamma(2)::EpGFP$ construct, in which the $\gamma(2)$ regulatory region drives GFP expression from the *endo16* basal promoter (Yuh and Davidson, 1996; Yuh et al., 1996), was produced by fusion PCR (Yon and Fried, 1989). The $\gamma(2)$ fragment was amplified from *SptbrBAC* (clone 31; J08 from arrayed library) using $\gamma(2)$ -Forward and $\gamma(2)$ -EpGFP-Reverse primers. EpGFP was amplified from the EpGFP expression vector (Cameron et al., 2004) using $\gamma(2)$ -EpGFP-Forward and GFP-Reverse primers. The fusion product was amplified using both fragments and the $\gamma(2)$ -Forward and GFP-Reverse primers. The resulting fragment was cloned into Promega pGEM-T Easy vector (Catalog #A1360) and fully sequenced to confirm proper fusion. The $\gamma(2)::EpGFP$ reporter construct was then amplified from the plasmid using $\gamma(2)$ -Forward and GFP-Reverse to produce linear fragments for injection.

$\gamma(2)$ -Forward: 5'-GTCTCTAGCAAGATATGTTACT-3'

$\gamma(2)$ -EpGFP-Reverse: 5'-ACAGTTTAACCCGGGAGATCTACTCTA-TAAACCACTACTGTACTCTA-3'

$\gamma(2)$ -EpGFP-Forward: 5'-TAGAGTACAGTAGTGGTTTATAGAGTA-GATCTCCCGGTTAAACTGT-3'

GFP-Reverse: 5'-ACTGGGTTGAAGGCTCTCAA-3'.

The Stratagene QuikChange Mutagenesis Kit (catalog #200518) was used to mutate or delete putative transcription factor binding sites on the $\gamma(2)::EpGFP$ plasmid. The resulting plasmids were sequenced to confirm introduction of the mutation. The following primer pairs were used to produce the *otxmut* $\gamma(2)::EpGFP$ and complex $\Delta\gamma(2)::EpGFP$ plasmids:

Otxmut $\gamma(2)::EpGFP$ -Forward: -CTGGTGATCGGTCAACTGATTCCTT-CCGGTTGGACGTGAA-3'

Otxmut $\gamma(2)::EpGFP$ -Reverse: 5'-TTCACGTCCAACCGGAAGGAAT-CAGTTGACCGATCACCAG-3'

Complex $\Delta\gamma(2)::EpGFP$ -Forward: 5'-TGTGCGTGCTTTACACCTGTC-TGGTGATCG-3'

Complex $\Delta\gamma(2)::EpGFP$ -Reverse: 5'-CGATCACCAGACAGGTG-TAAAGCAGCACA-3'

Complex $\Delta\gamma(2)::EpGFP$ -Reverse: 5'-CGATCACCAGACAGGTGTAAG-CACGACA-3'.

The mutated clones were checked by sequencing.

Ets and bHLH binding site mutations were introduced into $\gamma(2)::EpGFP$ by fusion PCR. “Left” fragments (produced using $\gamma(2)$ -Forward and the mutation's reverse primer) and “right” fragments (amplified with GFP-Reverse and the mutation's forward primer) were mixed to produce a megaprimer template for fusion PCR with $\gamma(2)$ -Forward and GFP-Reverse primers. The resulting full-length fragment was gel-purified and ligated into pGEM-T Easy vector for sequencing and amplification. The *etsmut1+2* $\gamma(2)::EpGFP$ construct was produced using *etsmut1* $\gamma(2)::EpGFP$ as a template for fusion PCR with the *etsmut2* mutation primers.

Etsmut1-Forward: 5'-GTCATTGACCTCAGATAGTCTGGTGATCG-3'
 Etsmut1-Reverse: 5'-TCACCAGACTATCTGAGGTCAATGACCGCTT-3'
 Etsmut2-Forward: 5'-CGAAGTTCACGTCCAAGAATCTGGGATTAGTT-3'
 Etsmut2-Reverse: 5'-GTCAACTAATCCAGATACCTGGACGTGAA-3'
 bHLHmut-Forward: 5'-CATTGACCTCTCTGCTACATGATCGGTCA-3'
 bHLHmut-Reverse: 5'-AGTTGACCGATCATGTAGCAGGAAGAGGT-3'.

The $\gamma(2)$ module was located by reiterative reporter assays as described (Smith and Davidson, 2008a). $\varepsilon\delta\gamma\beta\alpha::GFP$ was produced by fusion PCR between the 5' intergenic region of *tbr* (amplified from *Sptbr*BAC using *TbrA*-Forward and *TbrA*-Reverse primers) and GFP amplified with primers homologous to the basal promoter of *tbr* ($\varepsilon\delta\gamma\beta\alpha$ -GFP-Forward and GFP-Reverse). $\varepsilon\delta\gamma\beta\alpha::GFP$ was obtained by the same scheme using a different GFP forward primer, $\varepsilon\delta\gamma\beta\alpha$ -GFP-Forward. PCR fragments were cloned into pGEM-T Easy vector and sequenced. The $\varepsilon\delta\gamma\beta\alpha::GFP$, $\gamma\beta\alpha::GFP$, $\beta\alpha::GFP$, and $\alpha::GFP$ reporter constructs were produced from $\varepsilon\delta\gamma\beta\alpha::GFP$ using GFP-Reverse and the corresponding forward primers. $\gamma\alpha::GFP$, $\gamma::EpGFP$, and $\gamma(2)\alpha::GFP$ were generated by fusion PCR using an analogous method.

TbrA-Forward: 5'-GGAACGATACGAAACTTTG-3'
TbrA-Reverse: 5'-CTTAGGACCGTGTATATAC-3'
 $\varepsilon\delta\gamma\beta\alpha$ -Forward: 5'-CAGACAACCTGATTGCCTA-3'
 $\gamma\beta\alpha$ -Forward: 5'-TATAGGACCGTGTATATACCTC-3'
 $\beta\alpha$ -Forward: 5'-TATGTGTGCATGACTTTGCTT-3'
 α -Forward: 5'-AGATGGTTATTCTCCAGACTA-3'.

Shortened fragments of $\gamma(2)::EpGFP$ were produced by PCR amplification of $\gamma(2)::EpGFP$ using GFP-Reverse as a reverse primer and the following forward primers: $\gamma(2.1)$ -Forward, $\gamma(2.2)$ -Forward, $\gamma(2.3)$ -Forward, $\gamma(2.4)$ -Forward. These fragments were cloned into pGEM-T Easy vector and sequenced. The reporter fragment $\gamma(2.2-3)::EpGFP$ was produced by fusion PCR between the amplified region between the primers $\gamma(2.2)$ -Forward and $\gamma(2.3)$ -Forward (using $\gamma(2.2)$ -Forward and $\gamma(2.3)$ -Reverse) as well as amplified *EpGFP* with homology to $\gamma(2.3)$ produced by amplification with GFP-Reverse and $\gamma(2.3)$ -*EpGFP*-Forward.

$\gamma(2.1)$ -Forward: 5'-CATTTATTCGATCATCGA-3'
 $\gamma(2.2)$ -Forward: 5'-TGCTTTACAGTGATAACA-3'
 $\gamma(2.3)$ -Forward: 5'-TTGACGTGAACCTCGA-3'
 $\gamma(2.4)$ -Forward: 5'-CCATATAATTATAGATATATGA-3'
 $\gamma(2.3)$ -Reverse: 5'-GGGAGATCTACTCCGGAAGGGATTAG-3'
 $\gamma(2.3)$ -*EpGFP*-Forward: 5'-CCTTCCGGAGTAGATCTCCGGGT-3'.

Quantitative PCR

Embryos injected with recombiner BACs or reporter constructs were collected at the indicated time points. DNA and RNA were extracted using the Qiagen AllPrep DNA/RNA Mini kit (catalog #80204). Reverse transcriptase PCR was performed on the extracted RNA using the Biorad iScript cDNA synthesis kit (catalog #170-8890). BAC/reporter construct incorporation number and expression level were quantified by quantitative real-time PCR performed on extracted DNA and cDNA, respectively (Revilla-i-Domingo et al., 2004). The

single-copy gene *foxA* and two genes of well-characterized expression, *Spz12* (Wang et al., 1995) and *ubiquitin* (*ubq*; Oliveri et al., 2002; Ransick et al., 2002), were also quantified for comparison. The number of transcripts per embryo was determined by multiplying the fold difference in construct expression level (relative to *Spz12* or *ubq*) by the number of *Spz12* or *ubq* transcripts present at that time point, adjusting for GFP construct incorporation relative to *foxA* (Materna and Oliveri, 2008). *Spz12* and *ubq* standardizations gave consistent results; graphs shown are standardized relative to *Spz12*. The QPCR primers used are available online at: <http://sugp.caltech.edu/SUGP/resources/methods/q-pcr.php>.

Culture, microinjection, and fluorescence visualization

Culture and microinjection were performed as described (Flytzanis et al., 1985; McMahon et al., 1985) with the following modifications: eggs were not filtered prior to dejelling, and no BSA was added to dejellied eggs. Zygotes were injected with 10 pL of solution containing 150 molecules/pL of reporter construct or 40 molecules/pL of BAC and 120 mM KCl. HindIII fragment carrier DNA (4 nM) was added to injection solutions containing small reporter constructs. All BACs were linearized with *Ascl* prior to injection.

Translation and splice-blocking morpholino antisense oligonucleotides (MASO) were designed by GeneTools. For coinjections, MASO was added to the injection solution at the indicated concentrations. Embryos injected with a randomized mixture of morpholinos (IUPAC sequence: N₂₅) served as a mock-knockdown control.

Elk trans MASO: 5'-CGCTTCCGACATTGTGATGATTCTG-3'	400 μ M
Ets1/2 trans MASO: 5'-GAACAGTGCATAGACGCCATGATTG-3'	500 μ M
Ets4 splicing MASO: 5'-GCAAACTTCGCCAGTTGAGAACATG-3'	400 μ M
Erg trans MASO*: 5'-GCATATAACAAATTGAGGAACACTG-3'	200 μ M
Erg splicing MASO*: 5'-GGCCACTTCTGCAAAAACGAAC-3'	200 μ M
HesC trans MASO: 5'-GTTGGTATCCAGATGAAGTAAGCAT-3'	500 μ M
Tel trans MASO: 5'-CCTGTCTGTTAGAGCCGGTCCAT-3'	400 μ M

*Equal amounts of Erg trans and splicing MASOs were combined for Erg MASO injections (Oliveri et al., 2008).

pmar1 and *ets1/2* mRNA were obtained by plasmid transcription as described (Oliveri et al., 2002). Injection solution for mRNA coinjections contained 200 ng/ μ L *ets1/2* mRNA or 10 ng/ μ L *pmar1* mRNA. The final concentration of injected transcript did not exceed the maternal (for *ets1/2*; Rizzo et al., 2006) or early blastula (for *pmar1*; Oliveri et al., 2002) transcript number by more than tenfold, as recommended to maintain binding specificity (Materna and Oliveri, 2008).

GFP expression pattern was evaluated at the indicated time points on an epifluorescence Axioscope 2 Plus microscope (Zeiss, Hallbergmoos, Germany). Images were recorded with an AxioCam MRm (Zeiss) and fluorescence overlays produced in Adobe Photoshop CS 3.

Electrophoretic mobility gel shift assays and probe preparation

Gel shifts were performed using 12 h embryonic nuclear extract as described (Yuh et al., 1994). Double-stranded oligonucleotides were annealed and ³²P-labeled with Klenow DNA polymerase by the end-fill reaction. Underlined sequence represents overhang serving as a template for Klenow labeling.

H-Forward: 5'-GAGAGCCCTGTGCGTCTTACAGTGATAACAC-3'
 H-Reverse: 5'-GAGAGTGTATCACTGTAAAGCAGGCACAGGGGC-3'
 I-Forward: 5'-GAGAGTGTATCACTGTAAAGCAGGCAGGGTC-3'
 I-Reverse: 5'-GAGATGACCGCTTTGTGTTATCACTGTAAAGCAC-3'
 J-Forward: 5'-GAGAGTGATAACACAAAGCGGTCATTGACCTCTT-3'
 J-Reverse: 5'-GAGAAAGAGGTCAATGACCGCTTTGTGTTATCAC-3'
 K-Forward: 5'-GAGAAAGCGGTCATTGACCTCTTCTGTCTGGT-3'

K-Reverse: 5'-GAGAACCAGACAGGAAGAGGTCAATGACCGCTTT-3'
 L-Forward: 5'-GAGATTGACCTCTTCTGCTGCTGATCGGTCAA-3'
 L-Reverse: 5'-GAGATTGACCGATCACCAGACAGGAAGAGGTCAA-3'

M-Forward: 5'-GAGACCTGTCTGGTGATCGGTCAACTAATCCCTT-3'
 M-Reverse: 5'-GAGAAAGGGATTAGTTGACCGATCACCAGACAGG-3'
 N-Forward: 5'-GAGAGATCGGTCAACTAATCCCTTCCGGTTGGAC-3'

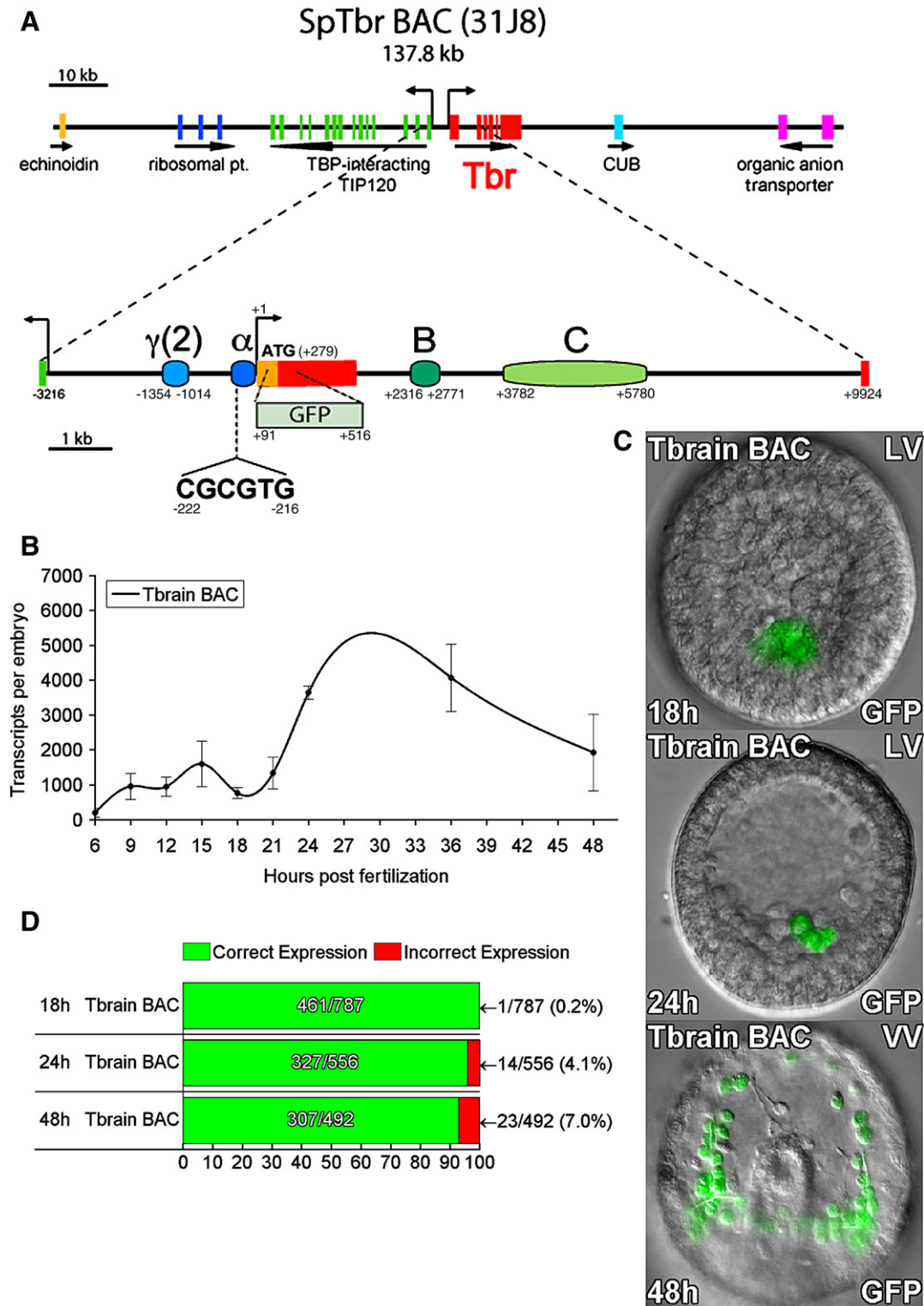


Fig. 1. Accurate skeletogenic expression of *tbr::GFP* BAC. (A) Map of the gene-rich *tbrain* BAC. The GFP coding sequence was recombined into *tbr* exon 1 as previously described (Damle et al., 2006). Positions of relevant *cis*-regulatory modules are indicated relative to the transcription start site (bent arrow) and exons (red boxes). (B) QPCR measurements of GFP mRNA in embryos bearing *tbr::GFP* BAC, 6–48 hpf. Transcript levels were normalized to measured BAC molecules in each sample, in this and all subsequent time-courses shown (see Materials and methods). Error bars indicate SEM in repetitions of the same experiments. (C) GFP fluorescence image overlays of *tbr::GFP* BAC-injected embryos at 18, 24, and 48 hpf. Expression is limited to the skeletogenic cells at all stages: LV, lateral view; VV, ventral view. (D) Summary of spatial expression statistics. Green and red bars indicate the fraction of embryos expressing GFP that showed fluorescence restricted to the SM cells (correct expression) vs. partially or completely ectopic fluorescence (incorrect expression), respectively. The total number of embryos injected is given in the denominators of these fractions.

N-Reverse: 5'-GAGAGTCCAACCGGAAGGGATTAGTTGACCGATC-3'
O-Forward: 5'-GAGACTAATCCCTTCCGGTTGGACGTGAACCTCG-3'
O-Reverse: 5'-GAGACGAAGTTCACGTCCAACCGGAAGGGATTAG-3'
P-Forward: 5'-GAGACCGGTTGGACGTGAACCTCGACCGCTGGTT-3'
P-Reverse: 5'-GAGAAACCAGCGGTGCAAGTTCACGTCCAACCGG-3'.

Results

Spatial and temporal expression pattern of recombineered *tbr::GFP* BAC

Abundant and ubiquitously distributed maternal transcript obscures the early zygotic expression pattern of the endogenous *tbr* gene. To visualize zygotic transcription we used a recombinant BAC, in which the coding region of GFP had been inserted at the start codon of *tbr* exon1 (Damle et al., 2006). Fig. 1B shows an expression time-course generated by quantifying GFP transcripts produced by this expression construct, *tbr::GFP* BAC, in embryos collected at 6–48 h after fertilization (hpf) and injection. GFP transcript number was normalized to the number of BAC DNA molecules incorporated per embryo. This was determined in QPCR measurements by comparing the incorporated genomic GFP coding sequence content to that of a known single copy gene, *foxa*.

Expression begins between 6 and 9 hpf, coincident with the disappearance of transcript encoding HesC, the predicted *tbr* repressor, from the micromeres between 8 and 12 hpf (Revilla-i-Domingo et al., 2007). There were ~1000 GFP transcripts/embryo between 9 and 21 hpf, increasing three-fold by 24 hpf, and remaining high at

36 and 48 hpf. This pattern of expression is consistent with previous time courses for endogenous *tbr* transcript (Oliveri et al., 2008 and additional unpublished data). The spatial expression pattern of *tbr::GFP* BAC was visualized in injected embryos by fluorescence microscopy at the blastula (18 hpf), mesenchyme blastula (24 hpf), and late gastrula (48 hpf) stages, as illustrated in Fig. 1C. Expression was highly specific to the SM lineage; the percentage of injected embryos displaying fluorescence anywhere else was ≤7% at all stages, and essentially zero at 18 h (Fig. 1D). The *tbr::GFP* BAC construct recapitulates both the spatial and temporal expression pattern of the endogenous gene with high fidelity.

The *tbr* gene is strikingly up-regulated by *pmar1* mRNA injection (Oliveri et al., 2002) and by *hesC* morpholino antisense oligonucleotide (MASO) injection (Revilla-i-Domingo et al., 2007), as required by the double-negative gate architecture. So indeed is the *tbr::GFP* BAC. Embryos coinjected with this construct and with *pmar1* mRNA, with *hesC* MASO, or with a random (N) MASO control were visualized by fluorescence microscopy at 18, 24, and 48 hpf. Both *pmar1* mRNA and *hesC* MASO injection resulted in increased amount of expression and grossly ectopic fluorescence relative to the control (Fig. S2A, C, E; Table S1). The *tbr::GFP* construct thus includes the genomic sequence required for these known regulatory inputs into the gene.

Ectopic GFP expression following HesC binding site mutation

A class C bHLH factor binding site (Iso et al., 2003) near the basal promoter is necessary for repression of a *Hemicentrotus tbr* construct

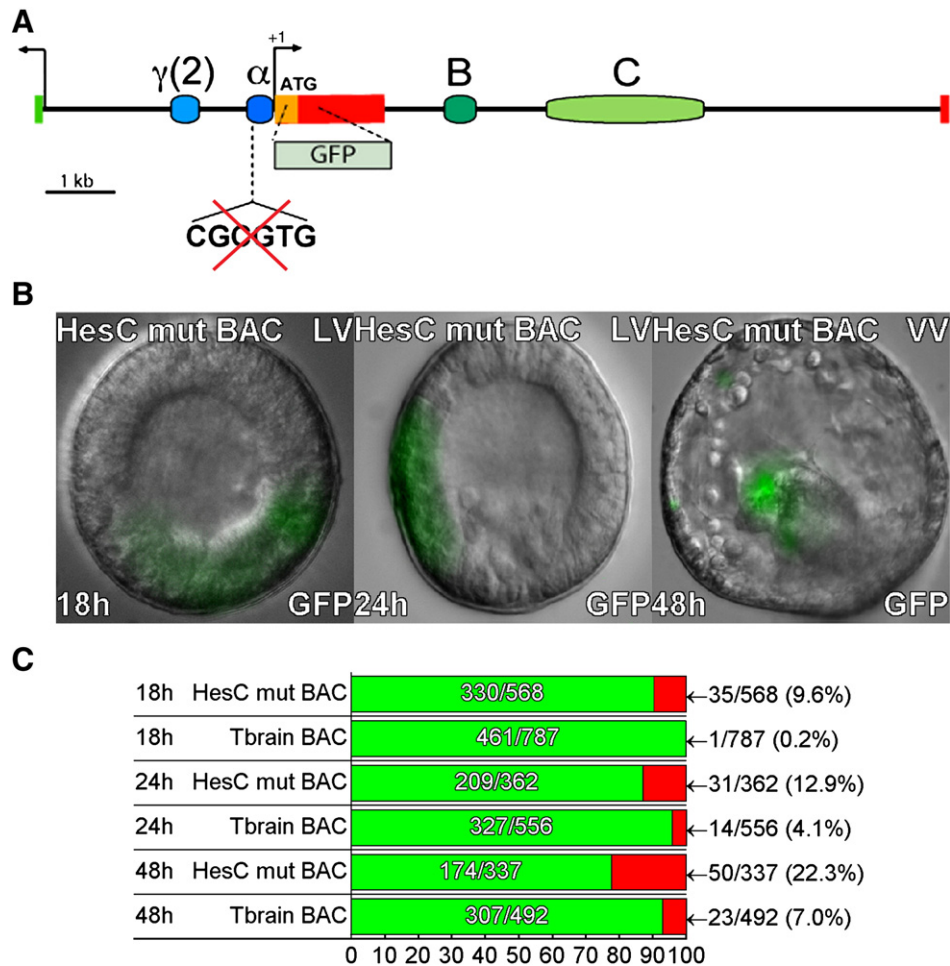


Fig. 2. Ectopic expression of *tbr::GFP* BAC following mutation of a HesC site. (A) Map of *tbr* locus. The location of the HesC binding site (CGCGTG) in the α region is indicated. (B) Examples of ectopic expression in ectoderm, endoderm, and NSM of *tbr::GFP* BAC in which this site had been mutated, at 18, 24, and 48 hpf. (C) Expression statistics for mutant and control BAC constructs as in Fig. 1D.

outside of the SM territory (Ochiai et al., 2008), and was thought to be a binding site for the HesC repressor implicated by gene network analysis in the control of *tbr* spatial expression in *S. purpuratus* (Oliveri et al., 2002; Revilla-i-Domingo et al., 2007). This sequence (CGCGTG) is conserved in the *S. purpuratus tbr* gene at -222 -217 relative to the transcription start site (Fig. 2A). To determine whether mutation of this single site would suffice to induce ectopic expression in the complete genomic context, a 4-bp mutation was introduced on the *tbr::GFP* BAC by means of homologous recombination. The mutation resulted in a significant increase in ectopic GFP expression relative to the *tbr::GFP* BAC control, while GFP expression in the SM lineage was unaffected (Figs. 2B, C). However, GFP misexpression was observed in only 10%, 13%, and 23% of embryos at 18 h blastula, 24 h mesenchyme blastula, and 48 h prism stages. This suggested that there could be additional undiscovered HesC sites; thus, by comparison, *pmar1* mRNA, which works by shutting down *hesC* expression, produced 49% ectopic expression by mesenchyme blastula stage, and the *hesC* MASO treatment used in these particular experiments 24% (Table S1). Computational analysis of the whole *tbr* regulatory

apparatus identifies several other potential HesC sites here not investigated; however, most of these lie in non-conserved regions of the sequence. Alternatively, this difference in misexpression rate caused by the mutation and that caused by *pmar1* mRNA and *hesC* MASO could be due to an indirect effect: both *pmar1* mRNA and *hesC* MASO injection cause the ectopic expression of *ets1/2*, an activator of *tbr* (see Discussion). In addition, we note that in a MASO injection the antisense oligo must be in excess to block the translation of the continuously transcribed *hesC*, which is not always attained, while the *pmar1* MOE produces enough transcriptional repressor to completely turn off the *hesC* gene.

Deletion of conserved intronic regions from *tbr::GFP* BAC

Ochiai et al. (2008) reported that Snail family consensus binding sites in a conserved intronic *cis*-regulatory module were necessary for repression of ectopic expression in a *Hemicentrotus tbr* reporter construct. The corresponding region, here identified as the B module (Fig. 3A), was deleted from the *S. purpuratus tbr::GFP* BAC by homo-

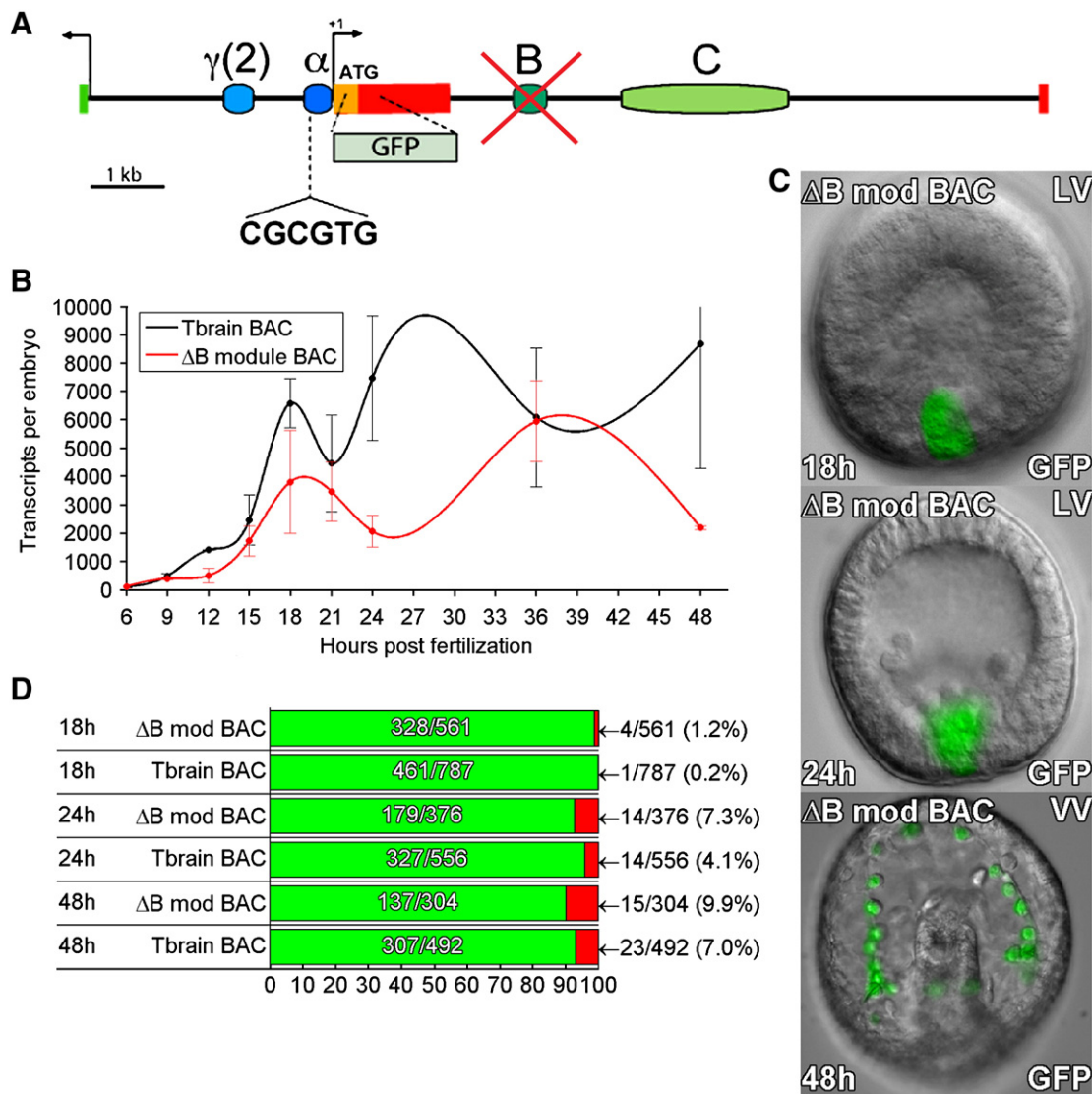


Fig. 3. Effects of deletion of B module from *tbr::GFP* BAC. (A) Map of *tbrain* locus. The B module was deleted by recombination (see Materials and methods). (B) QPCR time course of GFP mRNA levels generated by the B module deletion, and by the parental *tbr::GFP* BAC. In this and the following comparisons of diverse constructs data were obtained in experiments in which both constructs were injected into the same batches of embryos; i.e., the controls of each set of experiments were those of that comparison. Transcript levels were not adjusted for DNA incorporation rate of each construct. Error bars indicate SEM. No very large differences in expression level are observed in the B module deletion. (C) Examples of GFP fluorescence image overlays showing embryos expressing the B module deletion, at 18, 24, and 48 hpf. Expression is confined to SM cells. (D) Expression statistics for mutant and control BAC constructs as in Fig. 1D.

logous recombination. Quantification of GFP transcripts revealed no very significant differences in temporal expression pattern in the ΔB module BAC relative to the control, though there may be a transient depression of the level of activity soon after ingress (Fig. 3B). More importantly, there was no change whatsoever in the accuracy of expression caused by deletion of B module (Fig. 3C). Thus in *S. purpuratus*, the putative Snail binding site of B module has no detectable repressive spatial function when measured in complete genomic context.

An additional conserved region in the first intron of the *Tbrain* gene was identified as an enhancer in *Hemicentrotus* (Ochiai et al., 2008). When this region, here the C module, was deleted from the *tbr::GFP* BAC (Fig. 4A), a very significant decrease in GFP transcript levels was observed at all time points examined (Fig. 4B). Although the analogous deletion from a 7-kb *HpTbrain* reporter construct caused an increase in ectopic expression (Ochiai et al., 2008), we could detect no difference in the amount of ectopic expression produced by the ΔC module BAC vs. the control *tbr::GFP* BAC (Figs. 4C, D). Thus in *S. purpuratus*, C module in the context of the complete system appears to act as a quantitative enhancer of expression, but is not required for spatial accuracy of expression.

γ(2) module drives expression after ingress of the SM cells

A novel *cis*-regulatory module, *γ(2)*, which also mediates skeleto-genic expression, was identified in the 5' intergenic region of the *tbr* locus (Fig. 5A). It was found by means of iterative deletions from a large expression construct that included the whole intergenic region between *tbr* and the next gene upstream (Fig. S1). Successive deletions and results are shown in Fig. S3 and Table S2. To determine the function of *γ(2)* module in the context of the whole genomic regulatory system, this module was specifically deleted from the *tbr::GFP* BAC by homologous recombination. Study of the expression of this deletion construct revealed that it is expressed quite normally temporally and spatially until the time of ingress, but between 24 and 48 h a major decrease in expression levels is seen; this result is shown in Figs. 5B–D. In addition the *γ(2)* deletion produced a minor but significant increase in ectopic expression during this period, typically in the non-skeletogenic mesoderm. Thus in genomic context, *γ(2)* module functions after ingress. Since as shown in Fig. 4C module also functions during this period, we conclude that these two non-contiguous *cis*-regulatory modules collaborate in generating the definitive expression of the *tbr* gene in differentiated skeletogenic cells.

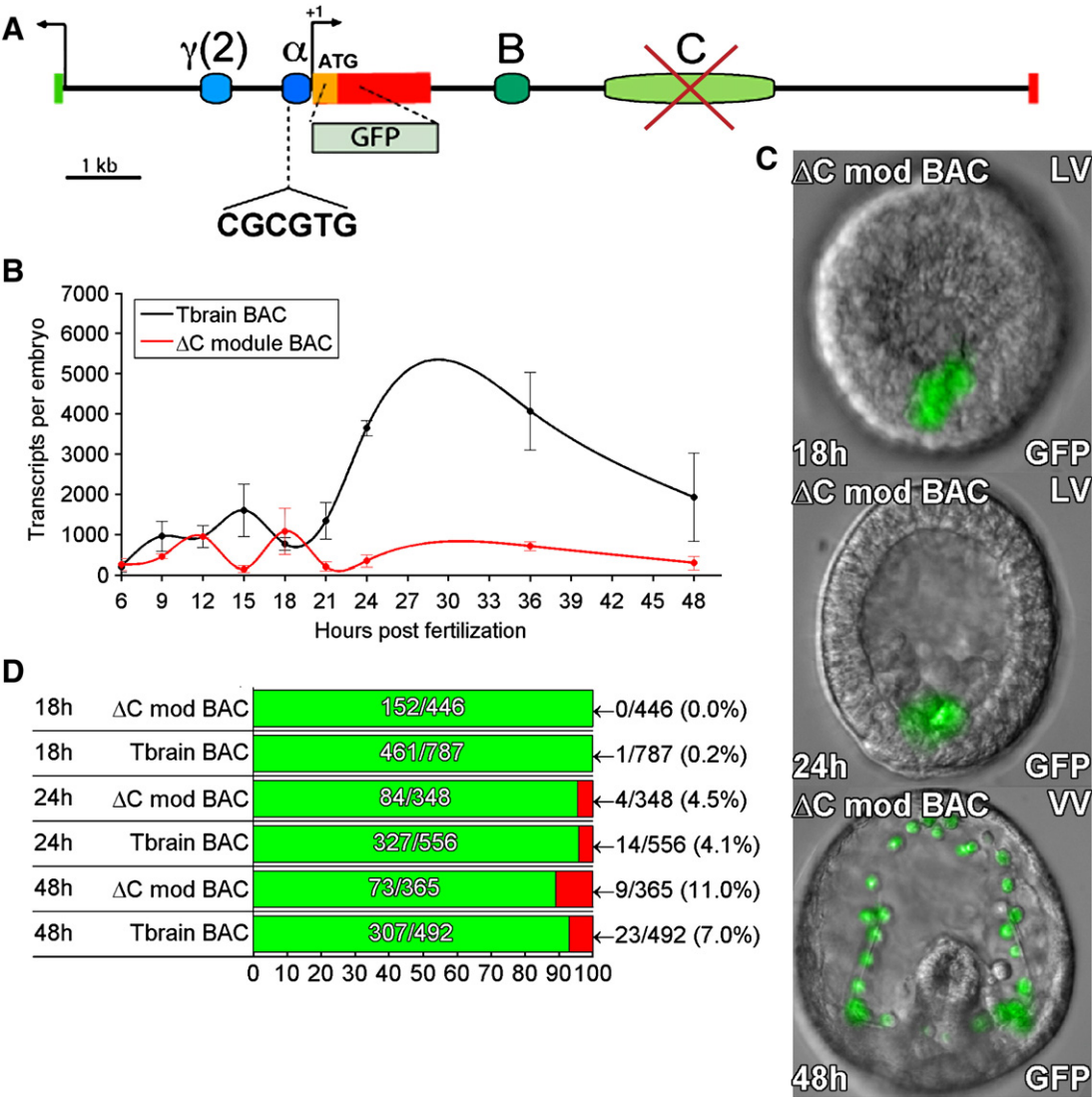


Fig. 4. Effects of deletion of C module from *tbr::GFP* BAC. (A) Map of *tbrain* locus. The C module was deleted by recombination (see Materials and methods). (B) QPCR time course of GFP mRNA levels generated by the C module deletion, and by the parental *tbr::GFP* BAC. Sharply decreased expression is observed relative to the control *tbr::GFP* BAC. Error bars indicate SEM of repeated experiments. (C) Examples of GFP fluorescence image overlays showing embryos expressing the B module deletion, at 18, 24, and 48 hpf. (D) Expression statistics for mutant and control BAC constructs as in Fig. 1D.

Expression of a short $\gamma(2)$ module construct lacking any other regulatory sequence

A standard minimal expression construct was created by fusing the $\gamma(2)$ module (Fig. 6 and Fig. S3) to the *endo16* basal promoter::GFP reporter (construct " $\gamma(2)::EpGFP$ "). On its own this basal promoter has no specific intrinsic spatial or temporal regulatory activity, but it mediates transcription in any domain of the embryo if provided with an exogenous *cis*-regulatory module active in that domain (Yuh and Davidson, 1996). In a head-to-head comparison the short $\gamma(2)::EpGFP$ construct is expressed just as accurately as is *tbr::GFP* BAC (Fig. 6B, C). We then compared the quantitative expression of this construct across developmental time to that of the *tbr::GFP* BAC from which $\gamma(2)$ module had been deleted, as for the experiments of Fig. 5. The simplest case we can consider is that the activity of the whole system is just the sum of the activities of its individual *cis*-regulatory modules. In this case the activity of the short construct should match the calculated difference between the activities of the *tbr::GFP* BAC and the *tbr::GFP* $\gamma(2)$ deletion BAC. This comparison is plotted in Fig. 6A.

There are two interesting aspects of the result. First, and most obviously, $\gamma(2)::EpGFP$ does not generate nearly as much activity per incorporated construct, in the period after 24 h, as is lost from the complete system when the $\gamma(2)$ module is deleted. To test whether this might be due to the exogenous *endo16* promoter used in this construct, we generated a construct in which the $\gamma(2)$ module was associated only with the endogenous *tbr* promoter, denoted in the maps shown in Fig. S3 as " α " (construct " $\gamma(2)\alpha::GFP$ "). This construct was expressed spatially with the same accuracy as $\gamma(2)::EpGFP$, and quantitatively at exactly the same level (Table S2; Fig. S3). Promoter strength or identity is therefore not the explanation for the weak expression per incorporated molecule of the short construct. There is some other reason, as discussed below, that the short construct functions far less efficiently in isolation than does the very same *cis*-regulatory module in context.

The second interesting aspect of the comparison in Fig. 6A is that in the period earlier than 21 h, the short construct is expressed at the same level, and also in the same skeletogenic cells as is *tbr::GFP* BAC. In other words, in the context of the whole system, Fig. 5B shows that $\gamma(2)$ module plays no role whatsoever prior to ingress, but in

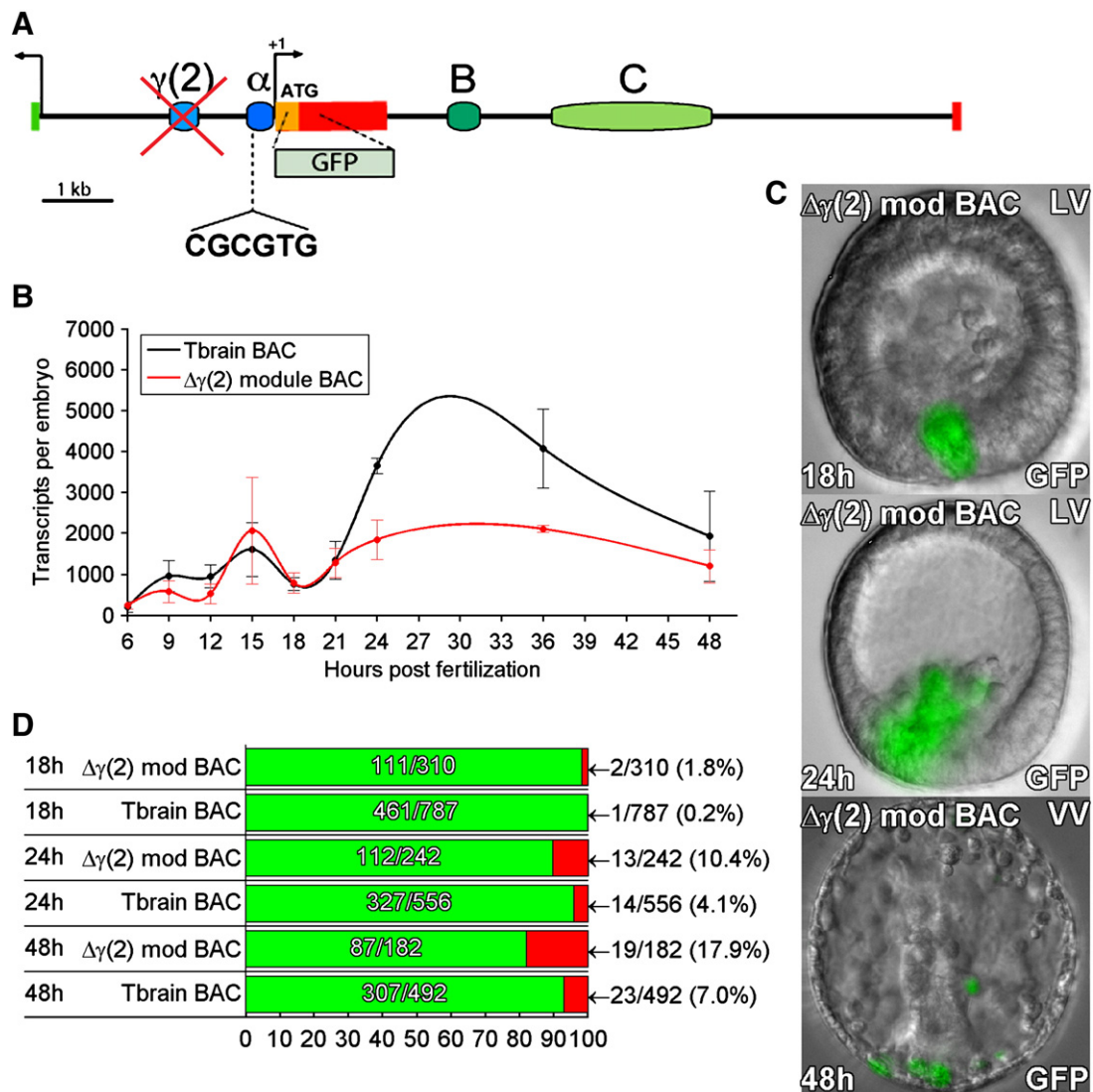


Fig. 5. Effects of deletion of $\gamma(2)$ module from *tbr::GFP* BAC. (A) Map of *tbrain* locus. The $\gamma(2)$ module was deleted by recombination (see Materials and methods). (B) QPCR time course of GFP mRNA levels generated by the $\gamma(2)$ module deletion, and by the parental *tbr::GFP* BAC. Significantly decreased expression is seen after 18 hpf. Error bars indicate SEM of repeated experiments. (C) Examples of GFP fluorescence image overlays showing embryos expressing the $\gamma(2)$ module deletion, at 18, 24, and 48 hpf. Increased ectopic GFP fluorescence was observed in the NSM cells of embryos bearing this deletion at 48 hpf, but not at 18 hpf. (D) Expression statistics for mutant and control BAC constructs as in Fig. 1D.

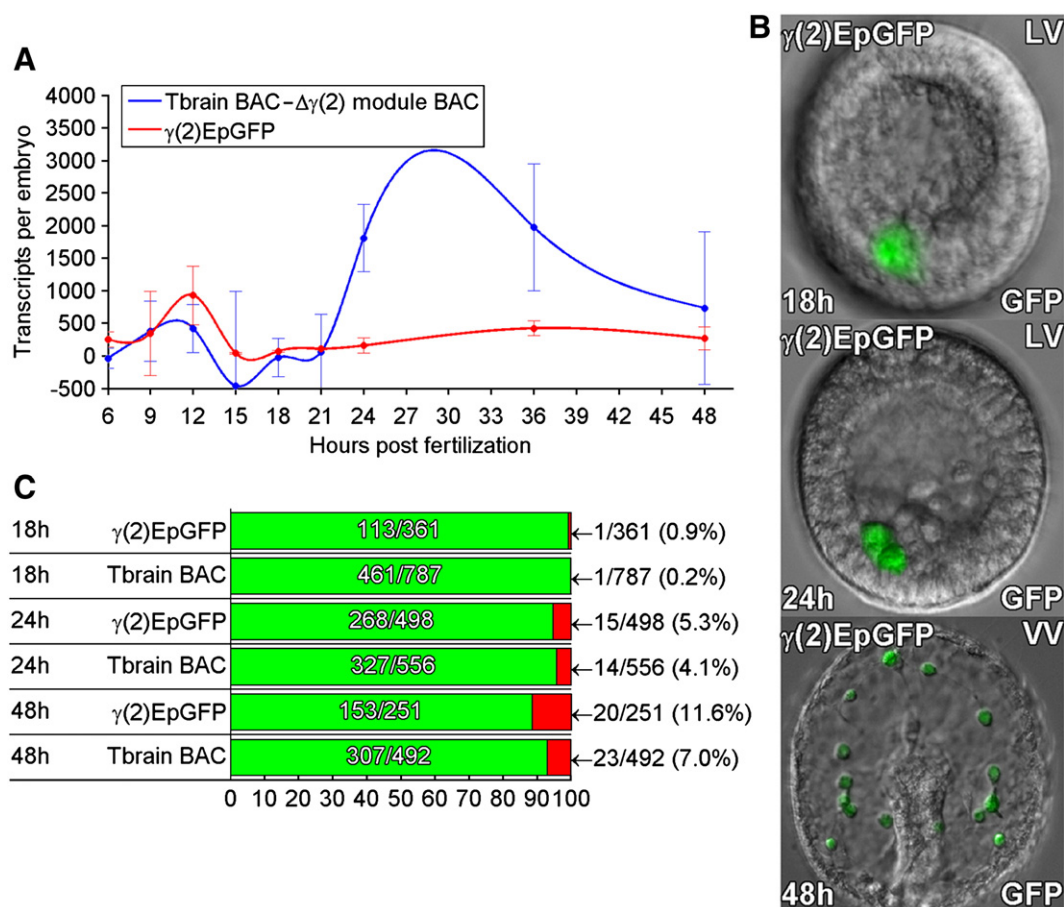


Fig. 6. Expression of a minimal $\gamma(2)$ module construct. A short construct consisting of the isolated $\gamma(2)$ module associated with our standard endo16-GFP expression vector was constructed as described in Materials and methods (construct $\gamma(2)::EpGFP$). (A) QPCR time course of GFP mRNA levels generated by $\gamma(2)::EpGFP$ (red) and co-plotted with the calculated difference between the time course produced by the control $tbr::GFP$ BAC and that produced by $tbr::GFP$ BAC from which $\gamma(2)$ module had been deleted (blue). All data are after normalization to the numbers of copies of the respective vectors incorporated at each time point, as above. Error bars indicate SEM. (B) Examples of GFP fluorescence image overlays showing embryos expressing $\gamma(2)::EpGFP$, at 18, 24, and 48 hpf. (C) Expression statistics for $\gamma(2)::EpGFP$ together with control $tbr::GFP$ BAC as in Fig. 1D.

isolation, as shown in Fig. 6A, it is capable of generating apparently normal spatial expression prior to ingress.

Given its accurate expression, we tested whether $\gamma(2)::EpGFP$ would respond similarly to $tbr::GFP$ BAC in perturbations of the upstream regulators. And indeed, injection of both *pmar1* mRNA and *hesC* MASO caused gross ectopic expression of the $\gamma(2)::EpGFP$ construct (Fig. S2; Table S1).

Ets family transcription factors regulate $\gamma(2)$ module

To identify the transcriptional activator(s) of the $\gamma(2)$ module and to determine whether *HesC* is a direct or indirect regulator, a gel shift analysis was carried out using nuclear extract from 12 hpf embryos. We found a 71-bp subregion of $\gamma(2)$ module (Fig. 7A) which drove GFP expression specifically in the SM, though less strongly than does the full $\gamma(2)$ module when incorporated in an expression construct ($\gamma(2.2-3)::EpGFP$; Fig. S3a; Table S3). As Fig. 7B shows, there are three putative kinds of DNA-protein complex in this region, which are found respectively in oligonucleotides containing Ets family consensus binding sites (Consales and Arnone, 2002), oligonucleotides containing an Otx family consensus binding site (Mao et al., 1994), and an oligonucleotide that included a 30-bp upstream region which produced an unresolved additional set of complexes. The activities of the $\gamma(2)::EpGFP$ construct and of derivatives in which each of these putative binding sites were mutated are given in the chart in Fig. 7C. Mutation of the putative Otx binding site had minor effect (from 38.4% in WT to 29.1% when mutated), while deletion of the 30-bp sequence

(which partially overlapped an Ets binding site) decreased the level of GFP expression and the number of injected embryos visibly expressing GFP. Mutation of either Ets binding site significantly reduced the number of GFP-expressing embryos, more strongly for site 1 than for site 2, and when both Ets binding sites were mutated, GFP expression was abolished. But none of these mutations produced any ectopic expression (e.g., Fig. S4a–g). Although no corresponding DNA–protein complex was observed, a consensus bHLH binding site in this region was also considered as a candidate *HesC* binding site. However, mutation of this site in $\gamma(2)::EpGFP$ affected neither quantitative nor ectopic expression (Fig. 7C; Fig. S4g).

There are five genes of the Ets family expressed in the SM by mesenchyme blastula stage, viz. *erg*, *ets1/2*, *ets4*, *elk*, and *tel* (Kurokawa et al., 1999; Rizzo et al., 2006). MASO directed against each of these Ets family members was co-injected with $\gamma(2)::EpGFP$. The results, also summarized in Fig. 7C, reveal that *Ets1/2* (and possibly *Elk*, which had a weak effect) are required for normal levels of expression of $\gamma(2)::EpGFP$. This raised the possibility that the spatial control of this short construct by *HesC* could be indirect, since the *ets1/2* gene is itself controlled by the *pmar1/hesC* double-negative gate. To test this, *ets1/2* mRNA was co-injected with $\gamma(2)::EpGFP$ or with $tbr::GFP$ BAC. There was a striking difference in the early expression (18 hpf) outcome: $\gamma(2)::EpGFP$ was now expressed ectopically all over the embryo but the $tbr::GFP$ BAC was not (Fig. S2g,h; Table S1). Thus the complete system encompassed in the $tbr::GFP$ BAC is subject to dominant repression by *HesC* as shown above, whereas the short construct is regulated only by *Ets1/2*. In

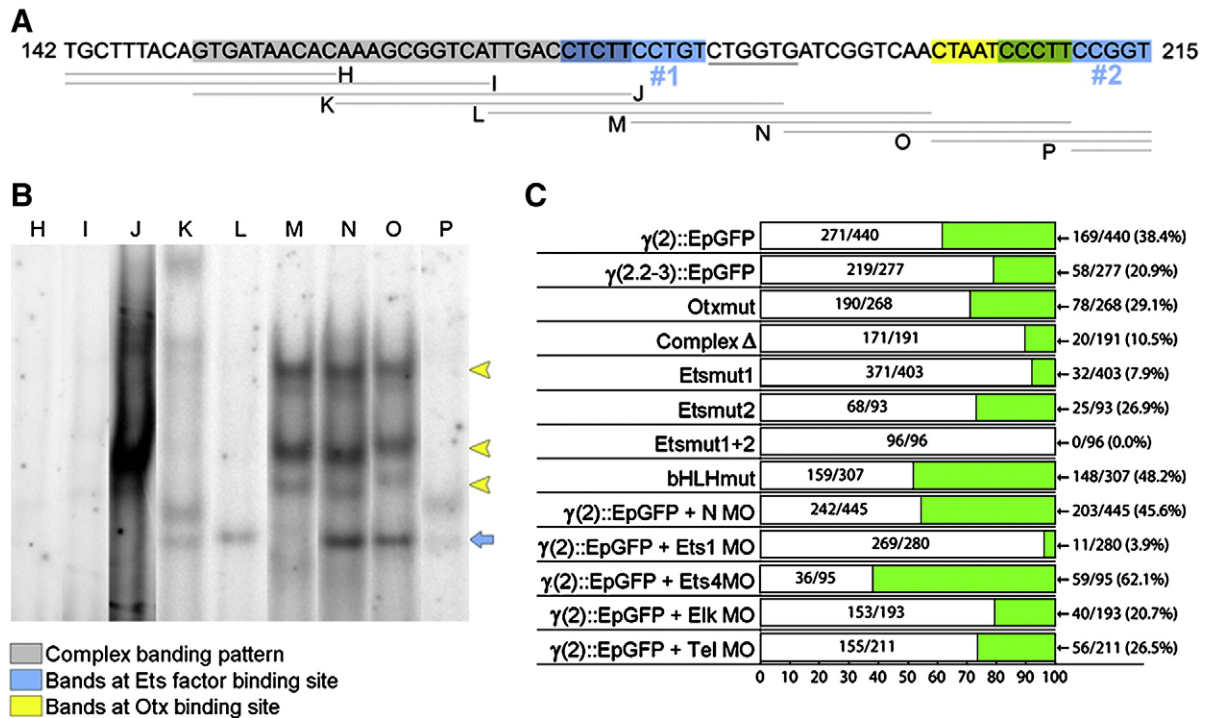


Fig. 7. Transcription factor binding sites in a subregion of $\gamma(2)$ module. (A) Map of a 71 bp subregion of $\gamma(2)$ module, showing the positions of oligonucleotides (H–P) used in the electrophoretic gel mobility shift assay. The inferred locations of putative target sites are indicated: Ets, blue; Otx, yellow; a possible bHLH site is underlined. (B) Gel shift results with oligonucleotides (H–P). Similar banding patterns are observed on oligonucleotides containing a putative Otx binding site (M–O, yellow arrowheads) and Ets factor binding sites (K, L and N, O, blue arrows). (C) Summary of expression results obtained with indicated site mutations and MASO treatments. Data are for 24 hpf embryos. All constructs, if they expressed at all, expressed accurately, and only quantitative expression results are shown. Embryos with ectopic GFP expression were rare and omitted from this figure for clarity; a full tally is provided in Table S3. Mutation of the putative Otx site had a minor effect and mutation of the bHLH sites had no effect on expression of $\gamma(2)::EpGFP$. However, mutation of either Ets binding site or deletion of the 30 bp region overlapping an Ets site (gray in (A)) dramatically decreased the number of embryos expressing GFP without affecting the spatial expression pattern. Mutation of both Ets sites eliminated almost all expression. Coinjection of $\gamma(2)::EpGFP$ with a random MASO (N MO) or MASOs directed against Ets family members Ets4 or Tel had no effect on expression, but *ets1/2* MASO and *elk* MASO dramatically decreased the number of embryos expressing GFP.

contrast, at later stages, when the $\gamma(2)$ module is functional, both *tbr::GFP* BAC and $\gamma(2)::EpGFP$ are ectopically expressed in *ets1/2* mRNA co-injection. This distinction in behavior excludes the possibility that $\gamma(2)$ module is literally redundant with the rest of the regulatory system.

An unexpected and important result of these MASO experiments was that introduction of *erg* MASO caused expansion of expression of both *tbr::GFP* BAC (Figs. 8A, D) and $\gamma(2)::EpGFP$ (Figs. 8C, D) into the NSM at 48 hpf. However, the *tbr::GFP* BAC construct from which the $\gamma(2)$ module had been deleted (Figs. 8A, B, D) was immune to this effect. Thus another late role of the $\gamma(2)$ module in the whole system is revealed: this function is to suppress transcription of the *tbr* gene in NSM in the gastrula stage embryo, a role necessitated by the expansion of *ets1/2* expression to the NSM by this stage.

Discussion

The *tbr* gene lies at an essential node, high in the gene regulatory network subcircuit which establishes the initial lineage specific regulatory state of the future skeletogenic mesoderm (SM; Oliveri et al., 2008). Network analysis predicts the key features of the genomic *cis*-regulatory code determining the transcriptional activity of this gene, and an initial motivation of this work was to explore these predictions. But it soon devolved that there are multiple components of this regulatory system: Ochiai et al (2008) identified several *cis*-regulatory modules in the *tbr* gene of a related species, while we had found a distinct *tbr cis*-regulatory module in a different region of the locus in *S. purpuratus*. Here we recount a system scale analysis that includes all known active modular units of the locus, based on recombinered BAC constructs which cover the complete locus and extend into the territories of the flanking genes on either side. The

network prediction that *tbr* is a primary target of the *pmar1-hesC* double-negative gate (Oliveri et al., 2002; 2008; Revilla-i-Domingo et al., 2007) was demonstrated true, and in this work we also solved the identity of the missing inferred control input that precludes *tbr* expression in the non-skeletogenic mesoderm (NSM). But in addition to resolving the functions of its various *cis*-regulatory inputs, we have gained unexpected insight into two other interesting aspects of the regulatory biology of the *tbr* gene. We discovered how different *tbr cis*-regulatory modules are deployed at different stages of development, and how, in this case, *cis*-regulatory inputs affect module choice. Not much is known about the subject of module choice, though it is obvious that the phenomenon is pervasive, as most regulatory genes appear to utilize multiple *cis*-regulatory modules (for review, Davidson, 2006). A related consequence, which has sharp implications for standard operating procedures in *cis*-regulatory analysis, was the demonstration that a “minimal enhancer” construct may display more functionality when introduced into an embryo than it actually executes in context, where what it does depends on whether it, rather than another module, is actually deployed. Finally the whole elaborate regulatory system we have revealed is cast into a particularly interesting light by the evolutionary novelty of this derived system, for as reviewed briefly in Introduction, only in echinoids is the *tbr* gene utilized at all in an embryonic SM cell lineage.

The early *tbr* control system

Disruption of the single HesC site in the α region of the *tbr::GFP* BAC produces a significant amount of ectopic expression in 18 and 24 hpf embryos, which though quantitatively minor is to be compared with the almost completely accurate expression of the parental BAC (Fig. 2, Table S1). A higher rate of ectopic expression was produced at

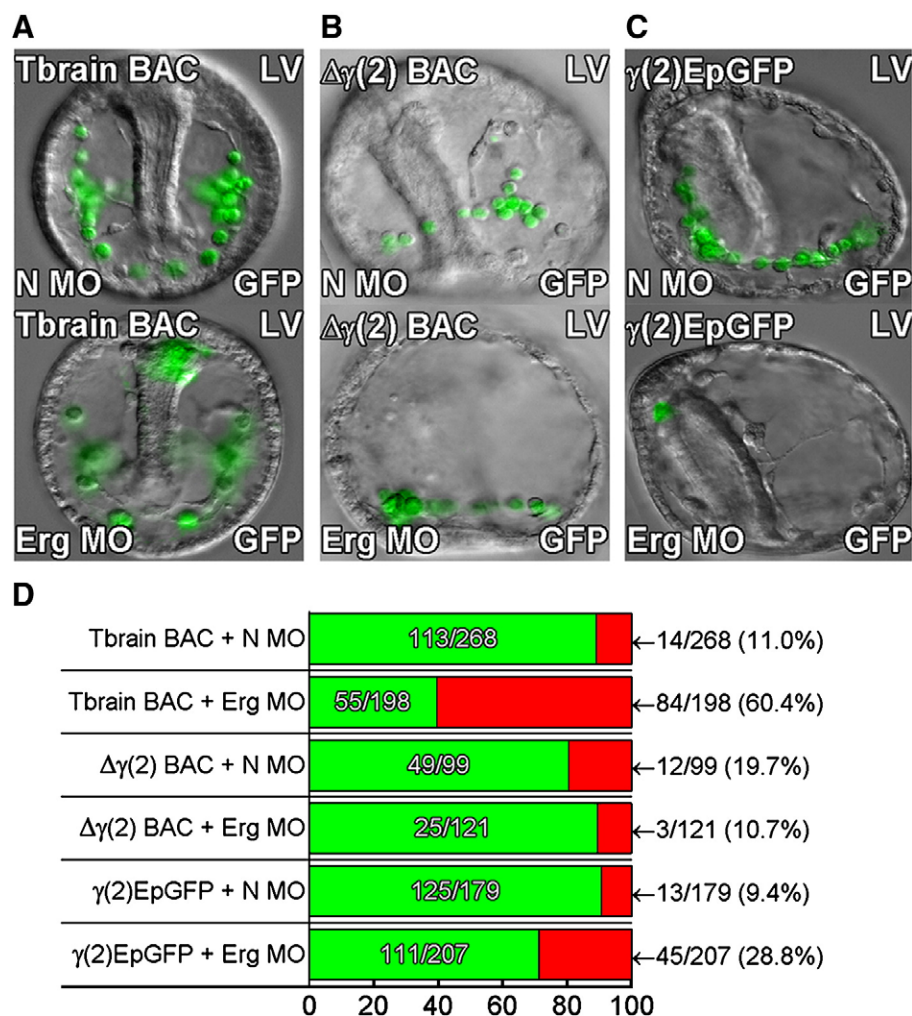


Fig. 8. Effect of *erg* MASO on expression of recombinant *tbr* BACs including and lacking $\gamma(2)$ module. (A) Examples of accurate spatial expression of control *tbr::GFP* BAC in presence of random MASO but ectopic expression in NSM cells in presence of *erg* MASO. (B) Example of accurate expression of *tbr::GFP* BAC from which the $\gamma(2)$ module had been deleted even in presence of *erg* MASO. (C) Effects of *erg* MASO similar to those in (A) are obtained with $\gamma(2)::EpGFP$. (D) Spatial expression statistics for these experiments.

these times by treatment with *hesC* MASO, using the wild type *tbr::GFP* BAC. The *hesC* MASO is clearly active as shown in earlier work (Revilla-i-Domingo et al., 2007), and as noted below, it sufficed in this study to produce 100% ectopic expression from the short $\gamma(2)::EpGFP$ construct later in development (Table S1). However, early in development when *hesC* is intensely transcribed everywhere in the embryo (except in the SM *pmar1* domain), it may be relatively difficult to block the presence of all HesC protein. We were mainly concerned to test in full genomic context the function of the single α module HesC site discovered by Ochiai et al (2008), and as noted above it is very possible that additional functional HesC sites exist elsewhere in the *tbr* locus.

The positive early control system consists of modules α plus C, as shown in the BAC deletions of Figs. 4 and 5. However, Figs. 4C, D shows that of these, C module is not required to produce accurate expression in the whole BAC. C module appears to contribute only a quantitative booster input since there is no increase in ectopic expression whatsoever when it is deleted, though there is a great decrease in level of expression (Fig. 4B). α module and its HesC site are able to do the job of ensuring that what expression remains is accurate. The location of any additional repressive HesC sites elsewhere in the locus would not have been tested in these deletions. Nonetheless, the significant destabilization of the very tight control executed by the early system operating in *tbr::GFP* BAC prior to ingress when the single known α module HesC site is destroyed, justifies the placement of this gene downstream of the *pmar1-hesC* double-negative gate.

tbr regulation after ingress

As shown very clearly in Fig. 5, when the upstream $\gamma(2)$ module is deleted from the complete system carried in *tbr::GFP* BAC, there is no effect of any kind on expression prior to ingress (21–22 hpf), either quantitative or spatial. But thereafter, the level of expression is greatly compromised; and in addition ectopic expression increases significantly, particularly in NSM cells (examples in Fig. 5C, 24 and 48 h embryos). The $\gamma(2)$ module is thus a late acting driver of expression in cells executing active skeletogenesis. It does not act alone, however, and again module C functions as a booster. These two modules interact cooperatively, since the sum of the expression in the late phase when C is deleted plus when $\gamma(2)$ is deleted does not equal the level of late expression when neither is deleted (Figs. 4 and 5).

The $\gamma(2)$ module has two different regulatory inputs, which probably use the same target sites. The experiments in Fig. 7 and Table S3 prove that the activating driver is indeed *Ets1/2*, interactions with which account entirely for its activity. We also demonstrated that the short $\gamma(2)$ module construct, $\gamma(2)::EpGFP$, responds sharply to *hesC* MASO; in fact by late gastrula this treatment causes 100% of embryos to mis-express the GFP reporter (Table S1). So also does global expression of the *Ets1/2* driver (Table S1). But $\gamma(2)$ module has no functional HesC site, and the effect of HesC on its expression is indirect. We can understand this at once by reference to the network architecture, for the *ets1/2* gene is itself a primary target of HesC repression immediately downstream of the *pmar1* double-negative

gate. Thus HesC MASO causes global ectopic expression of *Ets1/2* which in postgastrular embryos is normally confined to SM and NSM cells. That is why it causes global expression of $\gamma(2)::EpGFP$, the same effect on expression as direct injection into the egg of *ets1/2* mRNA (Table S1).

The experiments in Fig. 8 show that the reason the $\gamma(2)$ module does not express in the NSM even though the *Ets1/2* driver is present in these cells is that another NSM Ets family factor, Erg, acts to repress the activation potential of the module. After gastrulation *erg* is not transcribed in SM but continues to be expressed in NSM (Rizzo et al., 2006). Erg and Ets bind similar DNA target sites and so this is likely a case of competitive binding at the Ets sites, such that if the repressor Erg is present it wins. Thus *erg* MASO produces ectopic NSM expression of both the $\gamma(2)::EpGFP$ short construct and of *tbr::GFP* BAC (Fig. 8). But, the additional striking result in Fig. 8 is that *erg* MASO produces no ectopic NSM expression in the derivative of *tbr::GFP* from which $\gamma(2)$ module has been deleted. This reveals another late regulatory role of $\gamma(2)$ module in its normal context: not only does it cooperatively (with module C) drive expression in the SM, but it also represses it in the NSM.

Minimal module illusions, and the mechanism of $\gamma(2)$ module exclusion in early development

$\gamma(2)::EpGFP$ is a typical “minimal” expression construct, consisting of only the module itself and a promiscuous basal promoter-reporter apparatus. It gave near perfect expression both early and late (Fig. 6C), though as pointed out above, the short construct is quantitatively much less active per copy relative to its function in context. This could be due to the much greater flexibility of the longer DNA “arm” separating the module from the promoter in the normal context, allowing a greater variety of productive contact conformations, or to a greater tendency of the individual construct units to interfere with one another in the incorporated concatenate, or to titration of activators by the large number of short construct copies, or to a combination of these. The main point is not this, but the shocking discovery that in context the $\gamma(2)$ module apparently produces no output whatsoever prior to ingress, while when isolated in $\gamma(2)::EpGFP$ it does function prior to ingress. We see immediately that in the short construct, where there is no other option, the basal promoter will use whatever it can get, so to speak. The short construct does not exactly “lie” about $\gamma(2)$ module functionality; rather it “exaggerates”: only a part of what it displays may be utilized in context, because there is another layer of control, module choice. The fact that the complete system minus the $\gamma(2)$ module functions the same in early development as when $\gamma(2)$ module is present shows directly that $\gamma(2)$ module provides no significant input while the early α plus C module system is running. It operates differently, not redundantly with the α plus C module system, as shown by the strikingly different response to Ets overexpression in pre-ingress embryos. The interactions controlling $\gamma(2)$ module revealed in this study can also explain why it is silent in the early embryo.

In the pre-ingress SM we believe that the same thing happens to $\gamma(2)$ module as happens in the post-ingress NSM. As network analysis has shown (Oliveri et al., 2008), just downstream of the regulatory targets activated by the *pmar1-hesC* double-negative gate (i.e., *ets1/2*, *alx1*, *tel*, and *tbr*), a positive feedback subcircuit is activated by inputs from these primary responders. The first gene in this subcircuit is none other than *erg*. It receives an input from *tbr* itself as well as from *ets1/2*, then forges interactions with *hex* and *tgif*, including a feedback onto *erg* from *hex*. As we have seen, in the context of the whole system the $\gamma(2)$ module is dominantly repressed by Erg in the presence of *Ets1/2*, and so in the pre-ingress SM, once *erg* is turned on and kept on, $\gamma(2)$ module should be inactive. This is a case of short range repression (Gray et al., 1994) since the gene is not silenced, only the $\gamma(2)$ module. The circuitry, summarized in Fig. 9A, is

fascinating. Essentially, *tbr* expression is the cause of $\gamma(2)$ module repression, via the negative feedback from the *tbr* target *erg*. Or in other words the *tbr* gene itself ends up controlling which regulatory module will be deployed actively, and the exclusion of $\gamma(2)$ module activation potential is probably the cause of deployment of the α -C module system that operates in the early embryo rather than $\gamma(2)$ module. Later when *erg* expression is extinguished in the SM (for reasons not yet known, as this occurs later than our comprehensive network analysis so far extends), $\gamma(2)$ module is called into action, also in collaboration with C. The alternative conformations implied by these deployments are diagrammed in Fig. 9B. This is our preferred model, but it is also possible that an insulator contributes to silencing $\gamma(2)$ module in the complete construct, since we observed that interposition of a large stretch of upstream sequence in $\gamma(2)$ expression constructs prevents expression (Table S2; Fig. S3).

There are at least two possible reasons that the short $\gamma(2)::EpGFP$ construct does not respond to Erg repression in the early SM: first, the Ets activator may have a competitive advantage when its target sites are brought into immediate proximity of the basal transcription apparatus, forming a stable activation complex; second, the $\gamma(2)::EpGFP$ construct runs on an exogenous, promiscuous promoter from the *endo16* gene, and Erg repression may require elements of the endogenous promoter. As usual, negative results are subject to various interpretations, and it is what the $\gamma(2)::EpGFP$ construct does that is more informative than what it does not do.

Evolutionary considerations: How could all this have happened?

Almost all of the embryonic SM specification and differentiation gene regulatory network appears also to be utilized in the skeletogenic centers in which the initial spines and test plates of the adult body plan are constructed during mid-late larval life (Gao and Davidson, 2008). This includes the *ets1/2* and *alx* genes, as well as the triple feedback *erg*, *hex*, *tgif* subcircuit genes, and downstream regulators as well. Since the same apparatus is evidently deployed in the skeletogenesis centers of the sea star larva (which has no embryonic skeletogenic mesoderm lineage whatsoever), all of these genes appear to be components of a pleisiomorphic echinoderm skeletogenic network (Gao and Davidson, 2008). This network was evidently linked *in toto* into the embryonic specification system defining the micromere lineage in the evolutionary branch leading to the euechinoids, the modern sea urchins which display a precociously-ingressing skeletogenic micromere lineage. But none of this pertains to the *tbr* gene, because this gene is not part of the adult skeletogenic apparatus in either sea urchins or sea stars (Gao and Davidson, 2008). As reviewed in Introduction, *tbr* is expressed in the embryonic endoderm in other echinoderm classes and in euechinoid embryos exclusively in the SM.

The acquisition of *tbr* by the embryonic skeletogenic control apparatus of the euechinoids is a classic case of co-option, here seen directly at the network level. The switch away from its pleisiomorphic endodermal function may have had nothing to do directly with the *tbr* co-option process, since many regulatory genes participate in multiple developmental processes. There is some evidence that a key role of *tbr* in sea star embryonic specification, to provide a necessary feed into the *otx* gene, an essential endoderm regulator, has been supplanted by a different gene in the euechinoids, viz. *blimp1* (Hinman et al., 2007; Hinman et al., 2003). But this could have happened before, during or after *tbr* acquired its skeletogenic role. One essential step we can infer in the co-option process was placing *tbr* under control of the *pmar1-hesC* double-negative gate, as pointed out earlier (Gao and Davidson, 2008). This gate is not part of the adult skeletogenic apparatus either, and it is the definitive initiator of micromere specification. The other three first tier regulators answering to the double-negative gate also had to be placed under HesC control. Cis-regulatory studies on several double-negative gate targets (Smith and Davidson, 2008b) and

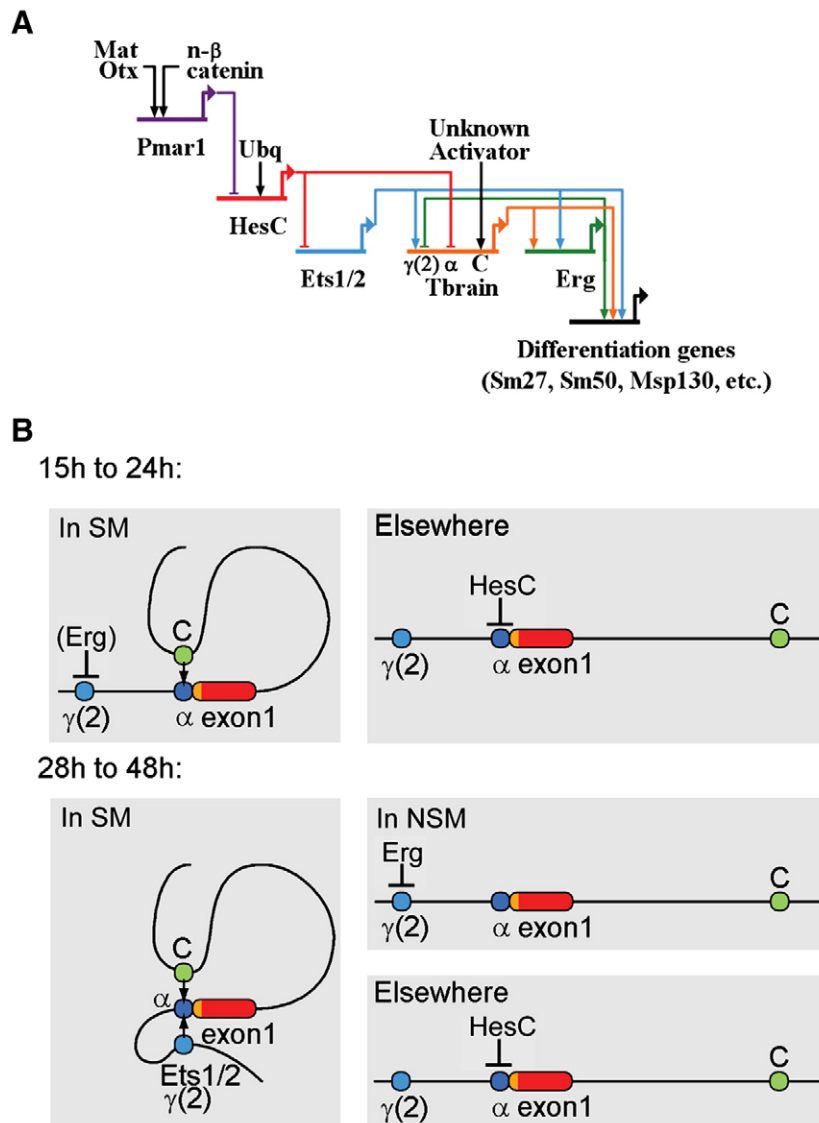


Fig. 9. Regulatory interactions of the *tbr* control system. (A) An updated *tbrain* regulatory subcircuit showing the inputs into the $\gamma(2)$ module from Ets1/2 and Erg identified in this study. The diagram displays interactions that occur in the SM prior to ingress. During this period the $\gamma(2)$ module is inactive (due to the negative feedback from the *erg* gene) and *tbr* expression is controlled by modules C and α , the site of HesC spatial repression (see text). Network diagram was constructed in BioTapestry (Longabaugh et al., 2009). (B) Schematics showing proposed interactions of *tbrain* cis-regulatory modules in different embryonic territories. This model is based on the now commonly assumed mechanism of interactions between distant cis-regulatory modules by DNA looping. Between 15 and about 21 hpf (Smith et al., 2008), dominant repression by HesC prevents *tbrain* expression outside of the skeletogenic micromere lineage. While Ets1/2 is present in the SM and then NSM during this period, *tbrain* activation in the PMCs occurs primarily through the C module due to the accumulation of the Erg repressor of $\gamma(2)$ module in the SM, as in (A). By ingress, HesC expression has disappeared in the NSM (Smith and Davidson, 2008b), and for unknown reasons expression of *erg* is extinguished in the SM cells; Erg and Ets1/2 are now both present in NSM but only Ets1/2 in the post-ingression SM. The result is that the $\gamma(2)$ module can now respond to SM Ets1/2 and contribute to *tbr* expression along with the module C booster, while $\gamma(2)$ module is shut off in the NSM by Erg and the expression of *tbr* elsewhere continues to be excluded by HesC.

unpublished data) show that one or two HesC sites does the job, and this aspect of the co-option process is easy to imagine.

But there is something special about *tbr* co-option, just because this gene is not part of the pleisiomorphic skeletogenic network apparatus, and the characteristics of $\gamma(2)$ module may hold the answer to the conundrum. The *tbr* gene has acquired several downstream targets in the SM, and so it is presumably useful as a differentiation driver. However unlike most others of these, *tbr* is never expressed in the NSM, as are *ets1/2*, *erg*, *hex*, etc. The reason, as we have seen, lies in the Erg repression function of the $\gamma(2)$ module. SM and NSM regulatory states greatly overlap but, because of $\gamma(2)$ module, *tbr* is an exception. In the evolutionary process leading to establishment of the embryonic euechinoid SM, $\gamma(2)$ module thus provided a mechanism for building a unique, non-skeletogenic mesodermal regulatory state. It is not the only one, for there is one other regulatory gene just downstream of the

double-negative gate that is also never expressed in the NSM, viz. *alx1*. The evolutionary role of $\gamma(2)$ module suggested here fits with its amazingly simple cis-regulatory construction, which depends essentially only on a couple of Ets1/2 target sites.

In summary, evolutionary co-option of *tbr* may have provided the special function of differentiating the SM from the NSM, just because the means of co-option included the appearance of $\gamma(2)$ module. Two other parts of this same function were provided by the still unknown mechanism by which transcription of the *erg* repressor is shut off in the SM, and by the equally SM-specific cis-regulatory apparatus of the *alx1* gene.

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Appendix A. Supplementary data

Supplementary data associated with this article can be found, in the online version, at doi:10.1016/j.ydbio.2009.08.005.

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